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NEWS 11 AUG 06 NEWS 12 AUG 13

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NEWS 22 SEP 24 NEWS 23 OCT 02

patents
EMBASE, EMBAL, and LEMBASE reloaded with enhancements
CA/CAplus enhanced with pre-1907 records from Chemisches
Zentralblatt

NEWS 24 OCT 19 BEILSTEIN updated with new compounds

NEWS EXPRESS 19 SEPTEMBER 2007: CURRENT MINDOWS VERSION IS V8.2, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.02c(JP), AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.

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13-14 14-15 15-16 16-17 3-4 4-5 5-6 12-13 12-17

1: CLASS 12: Atom 13: Atom 14: Atom 15: Atom 16: Atom 17: Atom 18: Atom 18: CLASS 12: Atom 13: Atom 14: Atom 15: Atom 16: Atom 17: Atom 18: CLASS 10:CLASS

STRUCTURE UPLOADED Ll

L1 HAS NO ANSWERS

Structure attributes must be viewed using STN Express query preparation.

SAMPLE SEARCH INITIATED 12:35:42 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 1514 TO ITERATE

100.0% PROCESSED 1514 ITERATIONS SEARCH TIME: 00.00.01

7 ANSWERS

168 ANSWERS

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*
PROJECTED ITERATIONS: 27946 TO 32614
PROJECTED ANSWERS: 7 TO 298

7 SEA SSS SAM L1

-> 8 11 888 full FULL SEARCH INITIATED 12:35:56 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 30126 TO ITERATE

100.0% PROCESSED 30126 ITERATIONS SEARCH TIME: 00.00.01

168 SEA SSS FUL L1

-> file caplus COST IN U.S. DOLLARS SINCE FILE FULL ESTIMATED COST

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FILE 'HOME' ENTERED AT 12:35:15 ON 02 NOV 2007

-> file reg

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STRUCTURE FILE UPDATES: 31 OCT 2007 HIGHEST RN 952181-70-3 DICTIONARY FILE UPDATES: 31 OCT 2007 HIGHEST RN 952181-70-3

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http://www.cas.org/support/stngen/stndoc/properties.html

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Chain nodes:
10 11 18
ring nodes:
1 2 3 4 5 6 7 8 9
Chain bonds:
8-10 10-11 10-18 11-12

5 6 7 8 9 12 13 14 15 16 17

ring bonds : 1-2 1-6 2-3 2-7 3-4 3-9 4-5 5-6 7-8 8-9 12-13 12-17 13-14 14-15 15-16 16-17

exact/norm bonds

8-9 10-11 10-18 11-12 exact bonds :

8-10 normalized bonds :

Robert Havlin

Robert Havlin

10/553,108

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FILE COVERS 1907 - 2 Nov 2007 VOL 147 ISS 20 FILE LAST UPDATED: 1 Nov 2007 (20071101/ED)

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http://www.cas.org/infopolicy.html

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20 L3

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L4 ANSMER 1 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 2007:1054426 CAPLUS Full-text
DOCUMENT NUMBER: 147:386026
INVENTOR(S): Preparation of nitrogenated heterocyclic derivatives as antagonists of chemokine receptor 5 (CCRS)
Kusuda, Shinya, Nishlyama, Toshihikov, Hashimura, Kazuya; Ueda, Junya, Shibayama, Shiro
Ono Pharmaceutical Co., Ltd., Japan
PCT Int. Appl., 185pp.
CODEN: PIXXD2
DOCUMENT TYPE: PAMILY ACC. NUM. COUNT: 1

PATENT	NO,		KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE	
		-		-									-		
WO 2007	105637		A1		2007	0920		WO 2	007-	JP54	684		21	0070	309
W:	AE, AG	, AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
	CN, CO	, CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
	GE, GH	, GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,
	KP, KR	, KZ,	LA,	LC,	LK,	LR,	LS,	LT.	LU,	LY,	MA,	MD,	MG,	MK,	MN,
	MW, MX	, MY,	MZ.	NA,	NG.	NI,	NO.	NZ.	OM,	PG.	PH,	PL.	PT.	RO,	RS,
	RU. SC	. SD.	SE.	SG.	SK.	SL,	SM,	sv.	SY,	TJ,	TM.	TN.	TR.	TT,	TZ,
	UA, UG	. us.	UZ.	VC.	VN.	ZA.	ZM.	ZW							
RW:	AT, BE								ES.	PI.	PR.	GB.	GR.	HU.	IR.
	IS, IT														
	BJ, CF														
	GH, GM														
	BY. KG														
PRIORITY APP	LN. INF	0.:						JP 2	- 000	6645	1		. 21	0060	310
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US COPYRIGHT 2007 ACS on STN 2007:671975 CAPLUS Pull-text L4 ANSWER 2 OF 20 CAPLUS ACCESSION NUMBER: 200

147:95554 Benzoxazole derivatives and related compounds as CETP inhibitors and their preparation, pharmaceutical composition and use for raising HDL and reducing LDL DOCUMENT NUMBER: TITLE:

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Robert Haylin
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Robert Haylin
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10/553,108
                                                                  2/40

cholesterol and treatment of atherosclerosis
Ali, Amjad; Hunt, Julianne A.; Kallashi, Florida,
Kowalchick, Jennifer E.; Kim, Dooseop, Smith, Cameron
J.; Sinclair, Peter J.; Sweis, Ramzi F.; Taylor, Gayle
E.; Thompson, Christopher F.; Chen, Liya; Quraishi,
                                                                                                                                                                                                                                                                              KG, KZ, MD,
PRIORITY APPLN. INFO.:
                                                                                                                                                                                                                                                                                                                                                RU, TJ, TM
                                                                                                                                                                                                                                                                                                                                                                                                   JP 2005-309642
JP 2006-76636
                                                                                                                                                                                                                                                                                                                                                                                                                                                        A 20051025
A 20060320
INVENTOR (5):
                                                                                                                                                                                                                                                                                                                                                     ARPAT 146:482079

THERE ARE 45 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
                                                                                                                                                                                                                                                                              OTHER SOURCE(S):
REFERENCE COUNT:
                                                                   Merck & Co., Inc., USA .
PATENT ASSIGNEE (S):
                                                                                                                                                                                                                                                                                                                                                LUS COPYRIGHT 2007 ACS on STN
2005:1026943 CAPLUS Full-text
                                                                   Merck & Co., Inc., USA
PCT Int. Appl., 294pp.
CODEN: PIXXD2
Patent
English
                                                                                                                                                                                                                                                                                L4 ANSWER 4 OF 20
ACCESSION NUMBER:
DOCUMENT TYPE:
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                                                                                                                                                                                                                                                                                                                                                 143:306325
                                                                                                                                                                                                                                                                                                                                                  143:306325
Substituted morpholine and thiomorpholine derivatives
 DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                                                                                                                                                                                                                                                TITLE:
                                                                                                                                                                                                                                                                                                                                                  as potassium channel openers, their preparation, pharmaceutical compositions, and use werzel Tornoe, Christian, Rottlaender, Mario; Khanzhin, Nikolay; Ritzen, Andreas; Matson, Will
                                                                                                                                                                                                                                                                               INVENTOR (9) .
             PATENT NO.
                                                                    KIND
                                                                                                                                                                                                                                                                                                                                                Khanzhin, Nikolay, Rit
Patrick
H. Lundbeck A/S, Den.
PCT Int. Appl., 88 pp.
CODEN: PIXXD2
Patent
English
                                                                                     20070621
                                                                                                                                                                           20061030
BZ, CA, CH,
FI, GB, GD,
KG, KM, KN,
MD, MG, MK,
PL, PT, RO,
TN, TR, TT.
             WO 2007070173
                                                                     A2
                                                                                                                                                                                                                                                                              PATENT ASSIGNEE(S):
SOURCE:
                                                                A2 20070621 MO 2006-US42208 20061030

MA, AT, AU, AZ, BA, BB, BG, BR, BM, BY, BZ, CA, CH,
CU, CZ, DE, DK, DM, DZ, EC, EZ, EG, ES, FI, GB, GD,
GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN,
LA, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK,
MY, MZ, NA, NG, NI, NO, NZ, OM, PC, PH, PL, PT, RO,
SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT,
US, UZ, VC, VN, ZA, ZM, ZM
CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, GM, GM,
MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, SY,
RU, TJ, TM
                      2007070173

M: AB, AG, AL,
CN, CO, CR,
GE, GH, GM,
KP, KR, KZ,
MN, MM, MX,
RS, RU, SC,
TZ, UA, UG,
RM: AT, BE, BG,
IS, IT, LT,
CP, CG, CI,
GM, KE, LS,
KG, KZ, MD,
APPLIN, INFO:
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LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                                                                                                                                                                                                                                                            PATENT NO.
                                                                                                                                                                                                                                                                                                                                                   KIND
                                                                                                                                                                                                                                                                                                                                                                   DATE
                                                                                                                                                                                                                                                                                                                                                                                                   APPLICATION NO
                                                                                                                                                                                                                                                                                                                                                                                                                                                                DATE
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                                                                                                                                                                                                                                                                                            WO 2005087754
 PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
                                                                                                                    US 2005-732168P
                                                                                                                                                                          P 20051031
                                                                   MARPAT 147:95654
L4 ANSWER 3 OF 20
ACCESSION NUMBER:
DOCUMENT NUMBER;
TITLE:
                                                        CAPLUS COPYRIGHT 2007 ACS on STN
2007:485607 CAPLUS <u>Pull-text</u>
146:482079
Preparation of 2-aminodihydrothiazine derivatives as
                                                                  Reparator, Araminor, Araminor, Araminor, Araminor, Araminor, Suzuki, Shinji, Sakaguchi, Gaku, Kato, Akira, Yukimasa, Akira, Hori, Akihiro, Koriyama, Yuji, Haraguchi, Hidekazu, Yasui, Ken, Kanda, Yasuhiko
                                                                                                                                                                                                                                                                                                                                                          INVENTOR (S):
                                                                   Haraguchi, Hidekazu, Y
Japan
PCT Int. Appl., 330pp.
CODEN: PIXXD2
Patent
Japanese
 PATENT ASSIGNEE (S):
 DOCUMENT TYPE:
 FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                                                                                                                                                                                                                                                             MX 2006PA10329
IN 2006CN03297
              PATENT NO.
                                                                    KIND
                                                                                     DATE
                                                                                                                      APPLICATION NO
                                                                                                                                                                                  DATE
                                                                                                                                                                                                                                                                                              NO 2006004599
             MO 2007049532 A1 20070503 W0 2006-JPJ21015 20061023
M1 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BM, BY, BZ, CA, CH,
CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
GE, GH, GM, GT, HN, HR, HU, JD, IL, TN, IS, JP, KE, KG, MK, KN,
KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LV, LV, MA, MD, MG, MK,
NN, MM, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO,
RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TT,
TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZM
RN, AT, BE, BG, CH, CY, CZ, DE, DK, EB, ES, FI, FR, GB, GR, HU, IE,
IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
CF, CG, CI, CM, GA, GR, GO, GM, ML, MR, ME, SN, TD, TG, GM, GH,
GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZM, AM, AZ, BY,
                                                                                                                                                                                                                                                                                PRIORITY APPLN. INFO.:
                                                                                                                                                                                                                                                                                                                                                                                                    US 2004-552574P
WO 2005-DK159
                                                                                                                                                                                                                                                                                                                                                                                                                                                                20040312
20050309
                                                                                                                                                                                                                                                                                                                                                   MARPAT 143:306325
5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
                                                                                                                                                                                                                                                                                OTHER SOURCE(S):
REFERENCE COUNT:
                                                                                                                                                                                                                                                                                 L4 ANSWER 5 OF 20 CAPLUS
ACCESSION NUMBER: 200
                                                                                                                                                                                                                                                                                                                                                           COPYRIGHT 2007 ACS on STN
                                                                                                                                                                                                                                                                                                                                                 LUS COPYRIGHT 2007 ACS on STN
2005:554644 CAPLUS <u>Full-text</u>
143:97280
Preparation of benzazepine derivatives as histamine H3
antagonists
Bailey, Nicholas; Bamford, Mark James, Dean, David
                                                                                                                                                                                                                                                                                TITLE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Robert Haylin
 10/553,108
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	Kenneth, Pickering, Paula Louise, Wilson, David	TD, TG
	Matthew; Witherington, Jason	AU 2004230367 A1 20041028 AU 2004-230367 20040413
PATENT ASSIGNEE (S) :	Glaxo Group Limited, UK	CA 2522266 A1 20041028 CA 2004-2522266 20040413
SOURCE:		EP 1614688 A1 20060111 EP 2004-727126 20040413
SOURCE:	PCT Int. Appl., 68 pp. CODEN: PIXXD2	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
DAGINENIE EVER		1E, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR
DOCUMENT TYPE:	Patent	
LANGUAGE:	English	
FAMILY ACC. NUM. COUNT:	1	PRIORITY APPLN, INFO.: JP 2003-109667 A 20030414
PATENT INFORMATION:		JP 2004-23032 A 20040130
		WO 2004-JP5237 W 20040413
PATENT NO.	KIND DATE APPLICATION NO. DATE	OTHER SOURCE(S): MARPAT 141:379923
		REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
WO 2005058837	A1 20050630 WO 2004-EP14380 20041215	RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
W: AE, AG, AL	, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,	
	, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,	L4 ANSWER 7 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN
GE, GH, GM	, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,	ACCESSION NUMBER: 2000:98525 CAPLUS Full-cext
LK, LR, LS	, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,	DOCUMENT NUMBER: 132:137396
NO. NZ. OM	, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,	TITLE: Phenylazole compounds, process for producing the same
	, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW	and drugs for hyperlipemia
	, KE, LS, MN, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,	INVENTOR(S): Umeda, Nobuhiro, Mochizuki, Nobuo, Uchida, Seiichi;
	, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,	Nishibe, Tadayuki, Yamada, Hirokazu, Ito, Kunihito;
	, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT,	Horikoshi, Hiromi
	, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,	PATENT ASSIGNEE(S): Nippon Soda Co., Ltd., Japan
MR, NE, SN		SOURCE: PCT Int. Appl. 92 pp.
		CODEN: PIXXD2
EP 1713778		DOCUMENT TYPE: Patent
	, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,	
	, LV, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK, HR, IS	LANGUAGE: Japanese
JP 2007514690	T . 20070607 JP 2006-544347 20041215	PAMILY ACC. NUM. COUNT: 1
US 2007060566	A1 20070315 US 2006-596503 · 20060615	PATENT INPORMATION:
PRIORITY APPLN. INFO.:	GB 2003-29214 A 20031217	
	WO 2004-EP14380 W 20041215	PATENT NO. KIND DATE APPLICATION NO. DATE
OTHER SOURCE(S):	MARPAT 143:97280	
REFERENCE COUNT:	6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS	' WO 2000006550 A1 20000210 WO 1999-JP4070 19990729
	RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT	W: AB, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
		DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS,
L4 ANSWER 6 OF 20 CA	PLUS COPYRIGHT 2007 ACS on STN	KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN,
ACCESSION NUMBER:	2004:902373 CAPLUS Full-text	MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM.
DOCUMENT NUMBER:	141:379923	TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD,
TITLE:	Preparation of phenylazole compounds as antioxidant	RU, TJ, TM
	drugs	RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
INVENTOR (S):	Mochiduki, Nobuo; Umeda, Nobuhiro; Uchida, Seiichi;	ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
	Ikeyama, Seiichi; Tsubokura, Shiro; Takada, Mitsumasa	CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PATENT ASSIGNEE(S):	Nippon Soda Co., Ltd., Japan	CA 2339123 A1 20000210 CA 1999-2339123 19990729
SOURCE:	PCT Int. Appl., 45 pp.	AU 9949297 Al 20000221 AU 1999-49297 19990729
	CODEN: PIXXD2	AU 753360 B2 20021017
DOCUMENT TYPE:	Patent	EP 1101759 A1 20010523 EP 1999-933152 19990729
LANGUAGE:	Japanese	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
FAMILY ACC. NUM. COUNT:		IE, SI, LT, LV, FI, RO
PATENT INFORMATION:	•	CN 1131217 B 20031217 CN 1999-809019 19990729
FAIRNI INFORMATION:		JP 2000290280 A 20001017 JP 1999-216581 19990730
PATENT NO.	KIND DATE APPLICATION NO. DATE	JP 2000281656 A 20001010 JP 1999-221789 19990804
PATENT NO.		JP 2000281658 A 20001010 JP 1999-221790 19990804
		OL 5000581038 W 50001010 OL 1222-551.20 12330804
NO 2004002162	A1 20041020 WO 2004 TRE227 2001011	HIS COLUMN DI CONTOLICO HIS CONTOLICO CONTOLICO
WO 2004092163	A1 20041028 WO 2004-JP5237 20040413	US 6342516 B1 20020129 US 2001-744786 20010126
W: AE, AG, AL	, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,	PRIORITY APPLN, INFO.: JP 1998-218316 A 19980731
W: AE, AG, AL CN, CO, CR	, AM, AT, AU, AZ, BA, BB, BG, BR, BM, BY, BZ, CA, CH, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD.	PRIORITY APPLN, 1NFO.: JP 1998-218316 A 19980731 JP 1998-222157 A 19980805
W: AE, AG, AL CN, CO, CR GE, GH, GM	, AM, AT, AU, AZ, BA, BB, BG, BR, BM, BY, BZ, CA, CH, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, HR, HU, ID, IL. IN, IS, JP, KE, KG, KP, KR, KZ, LC,	PRIORITY APPLN, INFO.: JP 1998-218316 A 19980731 JP 1998-22157 A 19980805 JP 1999-16846 A 19990126
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OCUMENT NUMBER:	128:57126			L4 ANSWER 11 OF 20 C	APLUS (	COPYRIGHT 20	7 ACS on STN		
ITLE:	Synthesis; cytotoxi	city, antitumor activi	y and	ACCESSION NUMBER:	1993:	570378 CAPL	JS <u>Full-text</u>		
		binding of two pyrazol		DOCUMENT NUMBER:	119:1				
		d to the antitumor age		TITLE:	Silve	r halide cole	or photographic photo	osensitive	
	and adozelesin	<b>-</b>			mater:	ials contain	ing two-equivalent ye	ellow couplers	
UTHOR (S):		nni; Cacciari, Barbara	Romagnoli,	INVENTOR (S):	Ikesu	, Satoru; Ki	a, Hiroshi, Kaneko,	Yutaka	
		ampiero, Gambari, Robe		PATENT ASSIGNEE (S):	Konica	Co., Japan			
		e, Marco; Ambrosino, P		SOURCE: .			Koho, 20 pp.		•
	Mongelli, Nicola, C	ozzi, Paolo, Geroni, C	ristina			: JKXXAF			
ORPORATE SOURCE:	Dipartimento di Sci	enze Farmaceutiche, Fe	rrara,	DOCUMENT TYPE:	Paten	t			
	I-44100, Italy			LANGUAGE:	Japan	ese		•	
OURCE:	Anti-Cancer Drug De	sign (1997), 12(7), 55	5-576	PAMILY ACC. NUM. COUNT:	1				
	CODEN: ACDDEA; ISSN	1: 0266-9536		PATENT INFORMATION:					
UBLISHER:	Oxford University P	ress							
OCUMENT TYPE:	Journal			PATENT NO.	KIND	DATE	APPLICATION NO	DATE	
ANGUAGE:	English								
EFERENCE COUNT:		CITED REFERENCES AVAIL		JP 04353844	A	19921208	JP 1991-153803	19910530	
	RECORD. ALL C	ITATIONS AVAILABLE IN	THE RE FORMAT	PRIORITY APPLN. INFO.:			JP 1991-153803	19910530	
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CCESSION NUMBER:	1996:466908 CAPLUS	Full-text		ACCESSION NUMBER:		6361 CAPLUS	rull-cexc		
OCUMENT NUMBER:	125:114620			DOCUMENT NUMBER:	114:6				
ITLE:		dazolylethyl)benzofura	n derivatives	TITLE:			tions of some new		
	as 5-lipoxygenase i			*			isoxazol-3-one deri	wat iven of	
NVENTOR (8):		Hachitani, Katsutoshi	; Nanbu,			ted biologic		vacives or	
	Pumio, Oonada, Shui						Abd El-Rahman, A. H.	. Dadeus D. C	
ATENT ASSIGNEE(S):	Ono Pharmaceutical			AUTHOR (S): CORPORATE SOURCE:			ra Univ., Mansoura,		
OURCE:	Jpn. Kokai Tokkyo K	ono, 120 pp.		SOURCE:			Chimie (1989), 34(9		
	CODEN: JKXXAF			SUORCE:			SN: 0035-3930	-10/, 1949-93	
OCUMENT TYPE:	Patent	•		DOCUMENT TYPE:	Journ		341 0033-3330		
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PATENT INFORMATION:	•			OTHER SOURCE(S):		ACT 114:6361			
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PATENT NO.		APPLICATION NO.	DATE	ACCESSION NUMBER:			US Full-text		
		JP 1994-270614	19941007	DOCUMENT NUMBER:	113:6		DS FUIT-CUAL		
JP 08109179		JP 1994-270614	19941007	TITLE:			dihydrobenzofuran he	rhicides	
RIORITY APPLN. INFO.:	MARPAT 125:114620	JF 1994-270614	19941007	INVENTOR (5):		e, Joseph E.			
THER SOUNCE(S):	MARPAT 125:114620	•		PATENT ASSIGNEE (S):			B, E. I., and Co., 'U	SA	
4 ANSWER 10 OF 20 CA	DI HE CODVETCHE 2007	ACC on CTM		SOURCE:			in-part of U.S. Se		
ACCESSION NUMBER:	1996:196719 CAPLUS			SOURCE .	aband		part or o,o, oo		
OCUMENT NUMBER:	124:261034	FULL-CEXT				: USXXAM			
TITLE:	Preparation and for	mulation of		DOCUMENT TYPE:	Paten				
1105.		alkylimidazoles and an	alogs As	LANGUAGE:	Engli				
		ents, antioxidants, an		FAMILY ACC. NUM. COUNT:					
	A2 synthetase inhib			PATENT INFORMATION:					
NVENTOR (S):		Hachitani, Katsutoshi	, Nanbu.						
	Pumio: Oonada, Shui			PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	• •
ATENT ASSIGNEE(S):	Ono Pharmaceutical								
OURCE:	Jpn. Kokai Tokkyo F			US 4881967	A	19891121	US 1988-202086	19880602	
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OCUMENT TYPE:	Patent			AU 8782152	A	19880616	AU 1987-82152	19871207	
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				CN 87107276	A	19881019	CN 1987-107276	19871207	
PATENT NO.	KIND DATE	APPLICATION NO.	DATE	CN 1021824	В	19930818			
			*******	DD 270532	A5	19890802	DD 1987-310042	19871207	
JP 07316150	A 19951205		19940524	US 4948418	A	19900814	US 1989-402178	19890830	
RIORITY APPLN. INFO.:		JP 1994-133575	19940524	US 5053071	A	19911001	US 1990-517892	19900502	
THER SOURCE(S):	MARPAT 124:261034			PRIORITY APPLN. INFO.:			US 1986-943365	A2 19861210	
							US 1988-202086	A3 19880602	
•									

				CN 87107276	A	19881019	CN 1987-107276	19871207	
PATENT NO.	KIND DATE APPLICATION NO.	DATE		CN 1021824	В	19930818			
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JP 07316150	A 19951205 JP 1994-133575	19940524		US 4948418	Α	19900814	US 1989-402178	19890830	
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							US 1988-202086	A3 19880602	
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	US 1989-402178	A3 19890830	3,000,17,34,1111	PATENT INFORMATION:					
OTHER SOURCE(S):	CASREACT 113:6148; MARPAT 113:6148			PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
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ACCESSION NUMBER: DOCUMENT NUMBER:	1989:534108 CAPLUS Full-text			PRIORITY APPLN. INFO.:	~1	1,000,02	DD 1985-276162	19850510	
TITLE:	Synthesis of some new benzodiazepine a	nd oxazenine		OTHER SOURCE(S):	CASREA	ACT 107:1154		13030310	
11106.	derivatives of expected biological act								
AUTHOR(S):	Habib, O. M. O., Abd El-Gawad, I. I.;			L4 ANSWER 17 OF 20 C					
CORPORATE SOURCE:	Fac. Sci., Mansoura Univ., Mansoura, E			ACCESSION NUMBER:			US Full-text		
SOURCE:	Polish Journal of Chemistry (1988), 62	(4-6), 543-7		DOCUMENT NUMBER:	91:81				
	CODEN: PJCHDQ; ISSN: 0137-5083			TITLE:	Develo	opment of ar	imagewise exposed c silver halide rec	ording material	
DOCUMENT TYPE: LANGUAGE:	Journal English			•	with a	priocographi	loper solution	orarna murarial	
LANGUAGE: OTHER SOURCE(S):	English CASREACT 111:134108			INVENTOR(S):			Kamitakahara, Atushi	: Mori, Keiichi	
OTHER SOURCE (S):				PATENT ASSIGNEE(8):	Konish	niroku Photo	Industry Co., Ltd.		
L4 ANSWER 15 OF 20 C	CAPLUS COPYRIGHT 2007 ACS on STN			SOURCE:	Ger. C	offen., 166	pp.	•	
ACCESSION NUMBER:	1988:21703 CAPLUS Full-text					: GWXXBX			
DOCUMENT NUMBER:	108:21703			DOCUMENT TYPE:	Patent				
TITLE:	Preparation of heterocyclic enol amide pharmaceuticals	derivatives as		LANGUAGE: FAMILY ACC. NUM. COUNT:	German	1			
PATENT ASSIGNEE(S):	Warner-Lambert Co., USA			PATENT INFORMATION:					
SOURCE:	Jpn. Kokai Tokkyo Koho, 78 pp.								
	CODEN: JKXXAF	•		PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
DOCUMENT TYPE:	Patent			DE 2823063	A1	19781130	DE 1978-2823063	19780526	
LANGUAGE: FAMILY ACC. NUM. COUNT:	Japanese .			DE 2823063	C2	19781130	DE 1978-2823063	13/80326	
PATENT INFORMATION:	•			JP 53146625	A	19781220	JP 1977-61917	19770526	
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PATENT NO.	KIND DATE APPLICATION NO.	DATE		US 4192680	A	19800311	US 1978-908913	19780524	
				PRIORITY APPLN. INFO.:			JP 1977-61917	A 19770526	
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ZA 8606973	A 19880427 2A 1986-6973 A 19870402 AU 1986-63285	19860912- 19860929		ACCESSION NUMBER: DOCUMENT NUMBER:	82:99		Full-text		
AU 8663285 AU 605747	B2 · 19910124	19060929		TITLE:			equivalent yellow o	ouplers	
DK 8504664	A 19870406 DK 1986-4664	19860930		INVENTOR (S):			W., Kirchhoff, Ger		
EP 221345	A1 19870513 EP 1986-113489	19861001		PATENT ASSIGNEE (S):		Sevaert AC			
R: AT, BE, CH	H, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE			SOURCE:	Ger. (	Offen., 17 p			
ES 2002398	A6 19880801 ES 1986-2338	19861001				: GWXXBX			
US 4921871	A 19900501 US 1987-121264	19871116		DOCUMENT TYPE:	Patent				
US 4874758	A 19891017 US 1988-164355	19880304		LANGUAGE: FAMILY ACC. NUM. COUNT:	German 1	n			
US 4868195	A 19890919 US 1988-165045 A 19890919 US 1988-166146	19880307 19880309		PATENT INFORMATION:	1				
US 4868200 US 4868199	A 19890919 US 1988-166146 A 19890919 US 1988-167264	19880309		TATELLY THE OLD ATTOM.					
US 4868205	A 19890919 US 1988-167272	19880311		PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
PRIORITY APPLN, INFO.:	US 1985-782623	A 19851001							
	US 1987-121264	A3 19871116		DE 2313989	Al	19740926	DE 1973-2313989	19730321	
OTHER SOURCE(S):	CASREACT 108:21703; MARPAT 108:21703		•	BE_812283	A2	19740916	BE 1974-1005793	19740314	
				CA 1016385	A1	19770830	CA 1974-195423	19740319 19740321	1
	CAPLUS COPYRIGHT 2007 ACS ON STN			GB 1434472	A	19760505	GB 1974-12564 DE 1973-2313989	A 19730321	
ACCESSION NUMBER: DOCUMENT NUMBER:	1987:515478 CAPLUS Full-text 107:115478			PRIORITY APPLN. INFO.:			75 13/3-2313989	. 13/30321	
TITLE:	2,3-Dihydrobenzo[b] furan derivatives			L4 ANSWER 19 OF 20 C	APLUS 4	COPYRIGHT 20	007 ACS on STN		
INVENTOR (S):	Boeckelmann, Juergen; Fanghaenel, Egon	ı; Grossmann,		ACCESSION NUMBER:	1974:	108307 CAPI	US Full-text		
	Norbert			DOCUMENT NUMBER:	80:10	8307			•
PATENT ASSIGNEE(S):	VEB Filmfabrik Wolfen, Fotochemisches	Kombinat, Ger.		TITLE:			tic acid (5,6-dimet)	oxycoumarone-2,3-	
	Dem. Rep.					boxylic acid	1)		
SOURCE:	Ger. (East), 4 pp.			AUTHOR (S):	Jha, C		laur Hair	ur tadir	•
DOCUMENT TYPE:	CODEN: GEXXAS			CORPORATE SOURCE: SOURCE:	Dep. C	cnem., Bhaga	alpur Univ., Bhagalı [ Chemistry (1973),	11/10) 989.90	
				SOURCE:			BSN: 0019-5103	1.01, 303-30	
LANGUAGE.	German								
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Robert Havlin
 L4 ANSWER 20 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1963:403485 CAPLUS Pull-text
  DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.;
                                                                                                                                           59:3485
                                                                                                                                           59:606b-h,607a-b
                                                                                                                                        59;606b-h,607a-b
2-Formyl-1,4-benzodioxane and 2-formyl-2,3-
dinydrobenzofuran
Migiti, Domenico, De Marchi, Franco, Rosnati, Vittorio
Ist. Super. Sanita, Rome
Gazzetta Chimica Italiana (1963), 93, 52-64
CODEN: GCITA9; ISSN: 0016-5603
Unavailable
  AUTHOR (S)
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    DOCUMENT TYPE:
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                               ANSWER 4 OF 20
    ACCESSION NUMBER:
    DOCUMENT NUMBER:
                                                                                                                                             Substituted morpholine and thiomorpholine derivatives
    TITLE:
                                                                                                                                           as potassium channel openers, their preparation, pharmaceutical compositions, and use wenzel Tornoe, Christian, Rottlaender, Mario, Khanzhin, Nikolay, Ritzen, Andreas, Watson, William Darrick
  INVENTOR (S):
                                                                                                                                          Khanzhin, Nikolay, Rit:
Patrick
H. Lundbeck A/S, Den.
PCT Int. Appl., 88 pp.
CODEN: PIXXD2
Patent
English
  PATENT ASSIGNEE(S):
SOURCE:
  DOCUMENT TYPE:
LANGUAGE:
    FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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A1 20050922 MO 2005-DX159

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                            MO 2005087754

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CN, CO, CR,
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LK, LR, LS,
NO, NZ, OM,
SY, TJ, TM,
RM: BM, GH, GM,
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EE, ES, FI,
RO, SE, SI,
MR, NE, SN,
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KR, KZ, LC,
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SK, SL, SM,
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A1 20061206 EP 2005-706619 20050309 .
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A 20070706 IN 2006-CN3297 20060912 .
A 20061208 NO 2006-4599 20061010
                               AU 2005221762
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EP 1727809
R: AT, BE, BG, IS, IT, LI,
CN 1930138
BR 2005008570
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	KIND DATE APPLICATION NO. DATE	
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	T 20070607 JP 2006-544347 20041215	
	A1 20070315 US 2006-596503 20060615	
IORITY APPLN, INFO.:		
	WO 2004-EP14380 W 20041215	
THER SOURCE(S): `		

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English

IN 2006CN03297 NO 2006004599

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Title compds. I [R1 = (un)substituted cycloalkyl: R2 = H, alkyl, cycloalkyl, etc.; X = a bond, CO, CO2, etc.; R3 = halo, alkoxy, CN, etc.; R4 = H, aryl, heteroaryl, etc.; n = 0-2 and their pharmaceutically acceptable salts, are prepared and disclosed as antagonists of histamine H3. Thus, e.g., II was prepared by reductive amination of N-(2,3,4,5-tetrahydro-IH-3-benzaepin-7-yl)4-morpholinecarboxamide (preparation given) with cyclobutanone. The activity of I was evaluated in the histamine H3 functional antagonist assay and it was revealed that numerous compds. of the invention possessed antagonism > 6.5 ptb. I as histamine H3 antagonists should prove useful in the treatment of neurol. disorders. Pharmaceutical compns. comprising I are disclosed.

annualizate
RL: PAC (Pharmacological activity), SPN (Synthetic preparation), THU
(Therapeutic use), BIOL (Biological study), PREP (Preparation), USES

Robert Haylin 10/553,108 PRIORITY APPLN, INFO.: 2004-412 20040312

MARPAT 143:306325

US 2004-552574P WO 2005-DK159

. STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT .

The invention relates to morpholine and thiomorpholine derivs. I, which are potassium channel openers. In compds. I, W is O or S; 2 is a bond or O, R1 is selected from halo. Cyano, C1-6 alkyl, C2-6 alkenyl, C2-6 alkynyl, C2-8 cycloalk(en)yl(oxy), etc., R2 is selected from halo, cyano, C1-6 alkyl, C2-8 cycloalk(en)yl(oxy), (un) substituted Ph, (un) substituted pyridinyl, etc., R3 is selected from C1-10 alkyl, C2-10 alkenyl, C2-10 alkynyl, C3-10 alkynyl, derivative III. The compds. of the invention express an EC50 value of less than 20  $\mu$ M, and in many cases less than 200 nM, in the assay of relative efflux through the KCNQ2

and in many cases less than 200 ins, in the assay of relative elifax through channel.

244540-15-2P, 2,3-Dihydrobenzofuran-2-carboxylic acid
(2,6-dimethyl-4-(morpholin-4-yl)phenyl)amide
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use), BIOL (Biological study); PREP (Preparation); USES
(Uses)
(drug candidate, preparation of substituted morpholine and thiomorpholine
derivs. as potassium channel openers)
864540-35-2 CAPLUS
2-Benzofurancarboxamide, N-[2,6-dimethyl-4-(4-morpholinyl)phenyl]-2,3dihydro- (CA INDEX NAME)

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

US COPYRIGHT 2007 ACS on STN 2005:564644 CAPLUS Full-text ANSWER 5 OF 20 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

Preparation of benzazepine derivatives as histamine H3 antagonists TITLE:

INVENTOR (S) .

antagonists
Bailey, Nicholas, Bamford, Mark James, Dean, David
Kenneth, Pickering, Paula Louise; Wilson, David
Matchew, Witherington, Jason
Glaxo Group Limited, UP
PCT Int. Appl., 68 pp.
CODEN, PIXXD2
Patent

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

Robert Havlin 10/553,108

16/40
(preparation of benzazepine derivs. as histamine H3 antagonists)
856904-12-6 CAPLUS
2-Benzofurancarboxamide, N-\_(3-cyclobutyl-2,3,4,5-tetrahydro-1H-3-benzazepin-7-yl)-2,3-dihydro-, mono(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

Robert Havlin

CRN 856904-11-5

CMF C23 H26 N2 O2

2 CM

76-05-1 C2 H F3 O2

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

US COPYRIGHT 2007 ACS on STN 2004:902373 CAPLUS Full-text ANSWER 6 OF 20 CAPLUS

141:379923

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

INVENTOR (S) :

141;379923
Preparation of phenylazole compounds as antioxidant drugs
Mochiduki, Nobuo, Umeda, Nobuhiro, Uchida, Seiichi, Ikeyama, Seiichi, Tsubokura, Shiro, Takada, Mitsumas Nippon Soda Co., Ltd., Japan
PCT Int. Appl., 45 pp.
CODEN, PIXXD2
Patent
Japanese PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC, NUM, COUNT: PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 2004092163	A1 20041028	WO 2004-JP5237	. 20040413
W: AE, AG, AL,	AM, AT, AU, AZ,	BA, BB, BG, BR, BW, BY	, BZ, CA, CH,
CN, CO, CR,	CU, CZ, DE, DK,	DM, DZ, EC, EB, EG, ES	, FI, GB, GD,
GE, GH, GM,	HR, HU, ID, IL,	IN, IS, JP, KE, KG, KI	P, KR, KZ, LC,
LK, LR, LS,	LT, LU, LV, MA,	MD, MG, MK, MN, MW, MX	K, MZ, NA, NI,
NO, NZ, OM,	PG, PH, PL, PT,	RO, RU, SC, SD, SE, SC	, SK, SL, SY,
TJ, TM, TN,	TR, TT, TZ, UA,	UG, US, UZ, VC, VN, YU	J, ZA, ZM, ZN
RW: BW, GH, GM,	KE, LS, MW, MZ,	SD, SL, SZ, TZ, UG, ZA	1, ZW, AM, AZ,

OTHER SOURCE(S):

MARPAT 141:379923

The title compds. I (wherein R = H or (un)substituted alkyl; A = (un)substituted imidazolyl or pyrazolyl; B = a bond or (un)substituted alkylene; Z = (un)substituted chroman-2-yl, 2,3-dinydrobenzofuran-2-yl, or 1,3-benzoxathlazol-2-yll or pharmaceutically acceptable salts thereof are prepared as antioxidant drugs. For example, the compound II was prepared in a multi-step synthesis. II showed antioxidant activity with ICSO of 3.3 µM in rat. I are useful for the treatment of kidney disorders, cerebrovascular disorders, retinal oxidation disorders, etc. (no data). Formulations containing I as an active ingredient were also described.

2-4163-52-69 764163-55-39 764163-55-19 784163-57-1P
784163-54-69 764163-65-1P
R1: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Uses) (drug candidate; preparation of phenylazole compds. as antioxidant drugs)
784183-52-6 CAPUS
2-Benzofurancarboxamide, 5-amino-2,3-dihydro-N-(4-(1H-imidazol-1yl)phenyl]-2,4,6,7-tetramethyl- (CA INDEX NAME)

784163-55-9 CAPLUS

2-Benzofurancarboxamide, 5-amino-2,3-dihydro-N-[3-(1H-imidazol-1-yl)phenyl]-2,4,6,7-tetramethyl- (CA INDEX NAME)

yl)phenyl)-2,4,6,7-tetramethyl-

10/553,108 Robert Haylin

784163-64-0 CAPLUS

2-Benzofurancarboxamide, 5-amino-2,3-dihydro-2,4,6,7-tetramethyl-N-[4-{lH-pyrazol-4-yl)phenyl}- (CA INDEX NAME)

784163-65-1 CAPLUS
2-Benzofurancarboxamide, 5-amino-2,3-dihydro-2,4,6,7-tetramethyl-N-[4-(2-methyl-1H-imidazol-1-yl)phenyl]- (CA INDEX NAME)

REFERENCE COUNT:

Robert Havlin

2-Benzofurancarboxamide, 5-amino-2,3-dihydro-N-[2-(1H-imidazol-1-yl)phenyl]-2,4,6,7-tetramethyl- (CA INDEX NAME)

784163-58-2 CAPLUS
2-Benzofurancetoxamide, 5-amino-2,3-dihydro-2,4,6,7-tetramethyl-N-[4-(1H-pyrazol-3-yliphenyl]- (CA INDEX NAME)

784163-61-7 CAPLUS

2-Benzo furancarboxamide, 5-amino-2,3-dihydro-2,4,6,7-tetramethyl-N-[3-(1H-pyrazol-3-yl)phenyl]- (CA INDEX NAME)

784163-62-8 CAPLUS
2-Benzofurancarboxamide, 5-amino-2,3-dihydro-2,4,6,7-tetramethyl-N-[4-(1-methyl-1H-pyrazol-5-yl)phenyl}- (CA INDEX NAME)

10/553,108 Robert Havlin

L4 ANSWER 7 OF 20 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

CAPLUS COPYRIGHT 2007 ACS on STN 2000;98525 CAPLUS <u>Full-text</u> 132:137396 Phenylazole compounds, process for producing the same

INVENTOR (S):

Verentylazota compounds, process for producing the sat and drugs for hyperlipenia Umeda, Nobuhiro, Mochizuki, Nobuo, Uchida, Seiichi, Nishibe, Tadayuki, Yamada, Hirokazu, Ito, Kunihito, Horikoshi, Hiromi Nippon Soda Co., Ltd., Japan PCT Int. Appl., 92 pp. CODEN: PIXXD2

APPLICATION NO.

WO 1999-JP4070

DATE

PATENT ASSIGNEE (S):

DOCUMENT TYPE:

Patent Japanese

FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO.

KIND DATE

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						•								<b>-</b>	-		
WO	2000	0065	50		A1		2000	0210	1	WO 1	999-	JP40	70		1	9990	729
	W:	ΑÉ,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,
		DE,	DK,	EE,	ES,	PI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,
		KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,
		MN.	MX,	NO.	NZ,	PL,	PT,	RO,	RU,	SD.	SE,	SG,	81,	SK,	SL,	TJ,	TM,
		TR.	TT.	UA.	UG,	US,	UZ,	VN,	YU,	ZA.	ZW,	AM,	AZ,	BY.	KG,	KZ.	MD,
		RU.	TJ.	TM													
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,	DK,
		ES.	FI.	FR.	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	BJ,	CF,	CG,
		CI.	CM.	GA.	GN,	GW,	ML.	MR,	NE.	SN.	TD,	TG					
CA	2339	123			Al		2000	0210		CA 1	999-	2339	123		1	9990	729
ΑU	9949	297			A1		2000	0221		AU 1	999-	4929	7		1	9990	729
ΑU	7533	60			B2		2002	1017									
EP	1101	759					2001			EP 1	999-	9331	52		1	9990	729
	R:	AT.					ES,										
					LV.			,					,				
CN	1131	217			В		2003	1217		CN 1	999-	8090	19		1	9990	729
JP	2000	2902						1017		JP 1	999-	2165	81		1	9990	730
	2000						2000			JP 1	999-	2217	89		1	9990	804
	2000						2000	1010		JP 1	999-	2217	90		1	9990	804
	6342									US 2	001-	7447	86			0010	
	APP											2183				9980	
										JP 1	998-	2221	57		1	9980	805
												1684				9990	
												1967				9990	
												2431				9990	

OTHER SOURCE(S):

PRI

MARPAT 132:137396

Phenylpyrazole and phenylimidazole compds. represented by general formula (I; wherein A represents (un) substituted imidazoly) or pyrazoly). B represents (un) substituted (CH2)% or (CH:CH)% Y = bond, O, S, 302, CO, COH2, Cl-5 alkyl-(un) substituted North (CH2)% or (CH:CH)% Y = bond, O, S, 302, CO, COH2, Cl-5 alkyl-(un) substituted North (CH2)% or (un) substituted and saturated or unsatd. heterocycle containing 1 to 4 N, O or S atoms, (un) substituted bargouptionyl or naphthoquinonyl) or pharmaceutically acceptable salts thereof are prepared Claimed are drugs for hyperlipemia which contain these compds. I as the active ingredient. Among all, compds. wherein Z is substituted throman-2-yl, e.g. dihydrobensofuran-2-yl, etc. have an effect of inhibiting the formation of lipid peroxides too. Thus, 6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid, 1-(4-aminophenyl)imidazole 4.0, 1-(3-dimethylaminopropyl)-3- ethylcarbodiimide hydrochloride 2.82, 1-hydroxybensotriazole 2.7g g, and 2.5 mL E23h were added to 30 mL DMA and stirred at room temperature for 20 h to give title compound (II). II and N-[4-(imidazol-1-yl)phenyl]-1-methyl-3- pyrrrolecarboxamide (III) at 25 mg/kg p.o. lowered total serum level of cholesterol 40 and 75%, resp., and serum triglyceride level by 62 and 91%, resp. A tablet formulation containing I was prepared 25660-78-JP 23660-84-JP 236650-84-JP 236650-54-JP 236650-78-JP 236650-78-JP 236650-78-JP 236650-78-JP 236650-78-JP 236650-78-JP 236650-78-JP 236650-78-JP 236650-78-JP 236650-78-JP

21/40

256660-84-1 CAPLUS

2-Benzofurancarboxamide, 5-(acetyloxy)-2,3-dihydro-N-[4-(1H-imidazol-1-yl)phenyl]-2,4,6,7-tetramethyl- (CA INDEX NAME)

10/553,108

23/40

Robert Havlin

REFERENCE COUNT:

THERE ARE 55 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 9 OF 20 CAPLUS ACCESSION NUMBER: 199 DOCUMENT NUMBER: 125

COPYRIGHT 2007 ACS on STN

1996:466908 CAPLUS Full-text 125:114620 .

TITLE

Preparation of (imidazolylethyl)benzofuran derivatives

INVENTOR (S):

Preparation of (imidazolylethyl)benzoturan derival as 5-lipoxygenase inhibitors
Hasegawa, Tomoyuki, Hachitani, Katsutoshi, Nanbu, Pumio, Oonada, Shuichi Ono Pharmaceutical Co. Japan
Jpn. Kokai Tokkyo Koho, 120 pp.
CODEN: JKXXAF

PATENT ASSIGNEE (S) : SOURCE:

DOCUMENT TYPE:

LANGUAGE -

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

JP 08109179
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
GI

DATE

APPLICATION NO.

19960430 MARPAT 125:114620

DATE

The title compds. [I, A = Cl-8 alkylene, B = 5-7-membered heterocycle containing 1-2 N atoms; G = OH, Cl-4 alkoxy, dialkylemino, etc., Rl, R2 = DE (wherein D = bond, Cl-8 alkylene, etc., E = OH, Cl-4 alkyl, cyano, alkoxycarbonyl, etc.); R4, R5 = H, Cl-8 alkyl, DE, etc.; n = 1-3], effective in treating and preventing thrombosis, atherosclerosis, etc., are prepared and formulated. Memylation of ethanol derivative II (R = OH) (preparation given) gave memylate II (R = MeSO3), which was heated with imidazole in toluene with stirring at 100° to give imidazole derivative II (R = 1-imidazoly)) (III). Hydrolysis of III with 4N HCl in MeOH gave diol salt IV, which showed 59% and 96% inhibition receiper LTMA and TUSA exerc.

Hydrolysis of III with 4N HCl in McOH gave diol sait IV, which showed 59% inhibition against LTM4 and TXB2, resp., at µM. 174956-03-20 RE. BAC (Biological activity or effector, except adverse), BSU (Biological study, unclassified), SPN (Synthetic preparation), THU (Therapeutic use), BIOL (Biological study); PREP (Preparation), USES (Uses) (preparation of (imidazolylethyl)benzofuran derivs, as 5-lipoxygenase

256661-23-1 CAPLUS

10/553,108

2-Benzofurancarboxamide, 2,3-dihydro-N-[4-(1H-imidazol-1-yl)phenyl]-2-methyl- (CA INDEX NAME)

L4 ANSWER 8 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1997:716591 CAPLUS Full-text DOCUMENT NUMBER: 128:57126

AUTHOR (S):

CORPORATE SOURCE:

DOCUMENT NUMBER:

TITLE:

128:57126
Synthesis, cytotoxicity, antitumor activity and sequence selective binding of two pyrazole analogs structurally related to the antitumor agents U-71,184 and adozelesin
Baraldi, Pier Glovanni, Cacciari, Barbara, Romagnoli, Romeo, Spalluto, Giampiero, Gambari, Roberto, Blanchi, Nicoletta; Passadore, Marco, Ambrosino, Piera, Mongelli, Micola, Cozi, Paolo, Geroni, Cristina Dipartimento di Scienze Parmaceutiche, Perrara, I-44100, Italy
Anti-Cancer Drug Design (1997), 12(7), 555-576
CODEN: ACDDRA; ISBN: 0266-9536
Oxford University Press
Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

English

UNGE: English
Two pyrazole analogs structurally related to the antitumor agents adozelesin and U-71,184
resp. were synthesized. By using a polymerase chain reaction approach, both compds, show
selective binding to A + T rich sequences exactly as reference compound U-71,184. In in
vitro assays, against Lill210 cell lines, both deriva, showed cytotoxicity in the pM range,
values comparable with the natural target compound (+)-CC-1065. The most active compound
showed very high antitumor activity in mice implanted with Lill210 cells (ILS% 363).
200264-86-4P

200264-86-4P
RI, RCT (Reactant), SPN (Synthetic preparation), PREP (Preparation), RACT (Reactant or reagent)
(preparation and antitumor activity and DNA binding of pyrazole analogs related to U-71,814 and adozelesin)
200264-86-4 CAPUS
HN-Indole-2-carboxylic acid, 5-[[(2,3-dihydro-2-benzofuranyl)carbonyl]amino]- (9CI) (CA INDEX NAME)

10/553,108

Robert Haylin

3.108 inhibitors)
174856-03-2 CAPLUS
2-Benzofurancarboxamide, 2,3-dihydro-5-hydroxy-7-[2-(1H-imidazol-1-yl)ethyl]-2,6-dimethyl-4-(1-methylethyl)-N-phenyl-, monohydrochloride
(9CI) (CA INDEX NAME)

174857-11-5F
RL, RCT (Reactant), SPN (Synthetic preparation), PREP (Preparation), RACT (Reactant or reagent)
(preparation of (imidazolylethyl)benzofuran derivs. as 5-lipoxygenase inhibitors)
174857-11-5 (APUS 2-Benzofurancarboxamide, 2,3-dihydro-7-(2-hydroxyethyl)-5-(methoxymethoxy)-2,6-dimethyl-4-(1-methylethyl)-N-phenyl- (9CI) (CA INDEX NAME)

L4 ANSMER 10 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER:
1996:196719 CAPLUS <u>Full-text</u>
124:26:1034
Preparation and formulation of dihydrobenzofuranylalkylimidazoles and analogs as antiinflammatory agents, antickvidants, and thromboxane A2 synthetase inhibitors
INVENTOR(S):
BASEGMAN, TOMODYAK!, Hachitani, Katsutoshi, Nanbu, Fumio, Oonada, Shuichi
PATENT ASSIGNEE(S):
ODCUMENT TYPE.
DOCUMENT TYPE.
LANGUAGE:
JAPANELS ACC. NUM. COUNT:
1

APPLICATION NO.

19951205

JP 1994-133575 JP 1994-133575

MARPAT 124:261034

The title compds. I  $\{R1, R2 = H, halo, etc.; A = alkylene, etc.; B = N-containing heterocyclic ring; R3 = H, acyl, etc.; R4 = H, alkyl, phenylalkyl, R5 = DE; D = alkylene, etc.; E = NR9R10, etc.; R3, R10 = H, alkyl, etc.; n = 1 - 3] are prepared. The title compound II.HCl was prepared in a multistep process starting from 2-(2-plus)olxyethyl)-3-methyl-4- acetyloxy-5-isopropyl-6-(2-methyl-2-propenyl)phenol. II.HCl in vitro at 10$ µM gave 96% inhibition of thromboxane B2 formation. 174856-03-2P

174856-03-29
RI. BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); TRU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of dihydrobenzofuranylalkylimidazoles and analogs as antiinflammatory agents, antioxidants, and thromboxane A2 synthetase

antinfiammeau, inhibitore)

174856-03-2 CAPLUS
2-Benzofurancarboxamide, 2,3-dihydro-5-hydroxy-7-[2-(1H-imidazol-1-yllethyl]-2,6-dimethyl-4-(1-methylethyl)-N-phenyl-, monohydrochloride
(9CI) (CA INDEX NAME)

174357-11-5P
RL: RCT (Reactant), SPN (Synthetic preparation), PREP (Preparation), RACT (Reactant or reagent) (preparation of dihydrobenzofuranylalkylimidazoles and analogs as antiinflammatory agents, antioxidants, and thromboxane A2 synthetase inhibitors)
174857-11-5 CAPLUS

27/40 Robert Haylin 10/553,108

L4 ANSHER 12 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER:
1991:6361 CAPLUS Full-text
114:6361
Synthesis and reactions of some new benzofurano[3,2-c]pyrazol-3-one and benzofurano[3,2-c]pyrazol-3-one and benzofurano[3,2-c]isoxazol-3-one and benzofurano[3,2-c]isoxazol-3-one derivatives of expected biological activity
Habib, O. M. O., Abd El-Rahman, A. H., Badawy, D. S.
CORPORATE SOURCE:
SOURCE:
Revue Roumaine de Chimie (1989), 34(9-10), 1949-55
CUDEN: RRCHAX; ISSN: 0035-3930
JOURNAL LANGUAGE:
OTHER SOURCE(S):
CASREACT 114:6361

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

Me salicylate condensed with α-chloroacetanilides to give the benzofurano-β-ketoanilides I (R = H, Cl, OMe). Treatment of I with N2H4.H2O, PhNNNH2, HONH2.HCl, polyphosphoric acid, and Mannich bases was studied. Reaction of the pyracolone derivative II with ClCH2COCl, PhN2+ Cl-, and Mannich bases was also investigated. 120948-2-0-6 Pl 130969-21-7P RL: RCT (Reactant) SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent) (preparation) FR (preparation and intramol. cyclocondensation of) 130968-20-6 CAPLUS

2-Benzofurancarboxamide, 2,3-dihydro-N-phenyl-3-(phenylhydrazono) - (9CI)

130968-21-7 CAPLUS 2-Benzofurancarboxamide, N-(4-chlorophenyl)-2,3-dihydro-3-

26/40 ZOPAU ZAPAU ZOPAU ZAPAU ZOPAU ZOPAU ZOPAU ZOPAU ZOPAU ZOPAU ZOPAU ZOPAU ZAPAU ZAPAU

L4 ANSWER 11 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1993:570378 CAPLUS Full-text DOCUMENT NUMBER: 119:170378

DOCUMENT NUMBER: TITLE:

119:170378
Silver halide color photographic photosensitive materials containing two-equivalent yellow couplers Ikesu, Satoru, Kita, Hiroshi, Kaneko, Yutaka Konica Co., Japan Jpn. Kokai Tokkyo Koho, 20 pp.
CODEN: JXXXAF

INVENTOR (9):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: Patent Japanese

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE APPLICATION NO DATE JP 04353844 PRIORITY APPLN. INFO.: JP 1991-153803 JP 1991-153803 19910530

The title photosensitive materials contain yellow couplers I (Z = nonmetallic atoms for forming 5-7-membered heterocyclic ring which may have substituents and condensed ring, R = substituent; n = 0-51. The invention produces photosensitive materials having good color rendition and provides high-quality color images having sufficient color d. and superior sharpness. 150004-08-3

IT RL: USES (Uses)

RN

(two-equivalent yellow photog. coupler)
150004-08-3 CAPLUS
Benzoic acid, 4-chloro-3-[{(2,3-dihydro-6-methoxy-3-oxo-2-benzofurany1)carbony1]amino}-, dodecyl ester (9CI) (CA INDEX NAME)

10/553,108 enylhydrazono) - (9CI) (CA INDEX

Robert Havlin

Robert Haylin

122529-46-8P 130968-18-2P 130968-19-3P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and reactions of)
122529-46-8 CAPUS
2-Benzagivanganhysmide 2 1-dibudge 2-avg Nachania (SCI) (CC TMP)

2-Benzofurancarboxamide, 2,3-dihydro-3-oxo-N-phenyl- (9CI) (CA INDEX

130968-18-2 CAPLUS

2-Benzofurancarboxamide, N-(4-chlorophenyl)-2,3-dihydro-3-oxo- (9CI) (CA INDEX NAME)

130968-19-3 CAPLUS

2-Benzofurancarboxamide, 2,3-dihydro-N-(4-methoxyphenyl)-3-oxo- (9CI) (CA

IT

130968-24-6P
RL: SPN (Synthetic preparation), PREP (Preparation)
(preparation of)
130968-24-0 CAPLUS
2-Benzofurancarboxamide, 2,3-dihydro-3-oxo-N-phenyl-2-(1piperidinylmethyl)-, monohydrochloride (9CI) (CA INDEX NAME)

L4 ANSWER 13 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 1990:406148 CAPLUS Full-text
DOCUMENT NUMBER: 111:6148
TITLE: Heterocyclic 2,3-dihydrobenzofuran herbicides

INVENTOR (S) Semple, Joseph E.

PATENT ASSIGNEE (S):

du Pont de Nemours, E. I., and Co., USA U.S., 57 pp. Cont.-in-part of U.S. Ser. No. 943,365, SOURCE:

CODEN: USXXAM Patent

English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DOCUMENT TYPE:

THE DAY THE CHARLES						
PATENT NO.	KIND	DATE	AP	PLICATION NO.		DATE
					-	
US 4881967	A	19891121	US	1988-202086		19880602
DK 8706415	A	19880611	DK	1987-6415		19871207
AU 8782152	A	19880616	ΑU	1987-82152		19871207
JP 63156787	A	19880629	JP	1987-307797		19871207
BR 8706588	A	19880726	BR	1987-6588		19871207
ZA 8709171	A	19880831	ZA	1987-9171		19871207
CN 87107276	A	19881019	CN	1987-107276		19871207
- CN 1021824	В	19930818				
DD 270532	A5	19890802	DD	1987-310042		19871207
US 4948418	A	19900814	US	1989-402178		19890830
US 5053071	A	19911001	US	1990-517892		19900502
PRIORITY APPLN, INFO,:			US	1986-943365	A2	19861210
			US	1988-202086	A3	19880602
			US	1989-402178	A3	19890830
OTHER SOURCE(S);	CASRE	ACT 113:6148;	MA	RPAT 113:6148		

3,108 31/40 Ro polyphosphoric acid gave benzodiazepines, e.g. II (X = NH), and benzoxazepines, (X = O). Benzodiazepines, e.g. II (X = NH), were obtained directly by the cyclocondensation of  $\beta$ -keto anilides, e.g. I (X = O), with o-(H2N)2C6H4. 127529-45-7 Robert Haylin 10/553,108

177529-45-9
RL: RCT (Reactant): RACT (Reactant or reagent)
(condensation reactions with o-phenylenediamine and o-aminophenol)
122529-46-8 CAPLUS
2-Benzofurancarboxamide, 2,3-dihydro-3-oxo-N-phenyl- (9CI) (CA INDEX NAME)

122529-49-1P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(preparation and cyclization of, benzyldiazepine derivative from)
122529-49-1 CAPLUS

2-Benzofurancarboxamide, 3-{(2-aminophenyl)imino}-2,3-dihydro-N-phenyl-(9CI) (CA INDEX NAME)

120519-55-9P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

Reactant or reagent (greparation and cyclization of, oxazepine derivative from) 122529-55-9 CAPLUS

2-Benzofurancarboxamide, 2,3-dihydro-3-[(2-hydroxyphenyl)imino]-N-phenyl-

(CA INDEX NAME)

ANSWER 15 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN SSION NUMBER: 1988:21703 CAPLUS <u>Pull-text</u> ACCESSION NUMBER: DOCUMENT NUMBER: 108:21703

Title compds. I (R = H, Cl. F, Cl-2 alkyl, Cl-3 alkoxy, Rl = H, Br, Cl, F, Me, MeO, cyano, F3C, F3CO, MF2CO, XI = O, S, R2 = H, Me, Et, R3 = H, Cl-4 haloalkyl, cyano, CoCl, M2C.CH, Mc, Cheoco, Cheoco, Mc, Certain, C, Carlo, C,

L4 ANSMER 14 OF 20 CAPLUS
ACCESSION NUMBER: 1989
DOCUMENT NUMBER:
TITLE:

PLUS COPYRIGHT 2007 ACS on STN
1989;534108 CAPLUS Full-text
111;134108
Synthesis of some new benzodiazepine and oxazepine
derivatives of expected biological activity
Habib, O. M. O., Abd Sl-Gawad, I. I., Badawy, D. S.
Fac. Sci., Mansoura Univ., Mansoura, Egypt
Polish Journal of Chemistry (1988), 62(4-6), 543-7
CODEN: PJCHDQ; ISSN: 0137-5083 AUTHOR (S) : CORPORATE SOURCE:

DOCUMENT TYPE:

Journal English LANGUAGE:

OTHER SOURCE (S): CASREACT 111:134108

Condensation of  $\beta$ -keto anilides, e.g. I (X = 0) with o-H2NC6H4X1H (X1 = NH, O), gave Schiff bases, e.g. I (X = NC6H4X1H-2). Cyclization of the Schiff bases in AcOH or

10/553,108 TITLE: Robert Havlin Preparation of heterocyclic enol amide derivatives as pharmaceuticals
Marner-Lambert Co., USA
Jpn. Kokai Tokkyo Koho, 78 pp.
CODEN: JKXXAF PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: Patent Japanese LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE JP 62081369 A 19870414 JP 1986-230231 19860930 US 4761424 19880802 US 1985-782623 19851001 ZA 8606973 AU 8663285 AU 605747 19880427 ZA 1986-6973 AU 1986-63285 19870402 19860929 19910124 DK 8604664 EP 221345 A 19870406 DK 1986-4664
A1 19870513 EP 1986-111449
DE, ES, FR, GB, GR, IT, LI, LU, NL, SE
A6 19880801 ES 1986-2338
A 19890501 US 1987-121264
A 19891017 US 1988-166355
A 19890919 US 1988-166146
A 19890919 US 1988-166146
A 19890919 US 1988-1672264
A 19890919 US 1988-167226
US 1985-782623
US 1985-782623
US 1985-782623
US 1987-7121264
CASREACT 108:21703; MARPAT 108:21703 19870406 DK 1986-4664 19860930 A Al EP 221345

R: AT, BE, CI
ES 2002398
US 4921871
US 4874758
US 4868195
US 4868200
US 4868200
US 4868199
US 4868205
PRIORITY APPLN. INFO.: 19861001 19861001 19861001 19871116 19880304 19880307 19880309 19880309 19880311 19851001 OTHER SOURCE(S):

The title compds. (I; Q = benzofuryl, benzothienyl, indolyl, benzopyranyl, benzothiopyranyl, etc., R5 = H, Cl-4 alkyl, alkoxy, C2-4 carbalkoxy, etc., R6 = C6-20 alkyl, styryl, etc., X = H, alkyl, m = 1, 2), useful as pharmaceuticals, are prepared A mixture of 0.085 mol furandione derivative II and 0.0749 mol antline derivite III in THF was stirred at room temperature under N, the solvent distilled in vacuo, and the solid product was refluxed in CH2Cl2 to give 85.2% enol amide IV. I showed ID50 against 5-lipoxygenase at 1.06-9.30M.

11926-22-8P 111926-23-9P 111926-28-4P

33/40

Robert Havlin

Robert Havlin

944-94-6P

111941-94-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological Study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological Study); PREP (Preparation), USES (Uses) (preparation of, as drug)
111926-22-8 CAPLUS
2-Benzofurancarboxamide, N-[4-[2-(3,4-dimethoxyphenyl)ethyl]phenyl]-2,3-dihydro-7-methoxy-3-oxo- (9CI) (CA INDEX NAME)

111926-23-9 CAPLUS
Naphtho[2,3-b] thran-2-carboxamide, N-[4-[2-(3,4-]
dimethoxypheny])ethyl]phenyl]-2,3-dlhydro-3-oxo- (9CI) (CA INDEX NAME)

111926-28-4 CAPLUS

Naphtho[1,2-b]furan-2-carboxamide, N-[4-[2-(3,4-dimethoxyphenyl]ethyl]phenyl]-2,3-dihydro-3-oxo-(9CI) (CA INDEX NAME)

CAPLUS

10/553,108

phtho[2,1-b]furan-2-carboxamide, 1,2-dihydro-N-[4-[2-(4-hydroxy-3-thoxyphenyl]ethyl]phenyl]-1-oxo- (9CI) (CA INDEX NAME)

DOCUMENT NUMBER: 91:81551
Development of an imagewise exposed light-sensitive color photographic silver halide recording material with a color developer solution Pushiki, Isamu, Kamitakahara, Atushi; Mori, Keiichi Konishiroku Photo Industry Co., Ltd., Japan INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: Ger. Offen., 166 pp. CODEN: GWXXBX DOCUMENT TYPE Patent German FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO KIND DATE APPLICATION NO DATE DE 2823063 DE 2823063 JP 53146625 JP 61023544 US 4192680 PRIORITY APPLN. INFO.: 19781130 19831103 19781220 19860606 19800311 A1 C2 DE 1978-2823063 19780526 JP 1977-61917

35/40

C5H11-tert VHCO (CH2) 30. -C5H11-tert

Yellow couplers having the formulas RCOCHRICONHR2 (R \* alkyl, cycloalkyl, or aryl; R1 \* a heterocycle; R2 \* aryl or a heterocycle), I (R3 \* alkyl, cycloalkyl, aryl, or a heterocycle; R, X1 \* 0 or NR5 where R5 \* alkyl, aryl, or acyl, and X and X1 are not the same; X2 \* 0, \$9, \$1 II (R6 \* alkyl, cycloalkyl, aryl, or a heterocycle; R7 \* H or R3 above; X3 \* C0, C5, 0, 8, alkylene, arylene, or a divalent heterocycle; X4 \* 0, S, acylimino, or sulfonylmino), and III (R8 \* aryl; R9 \* a noncleavable group; R10 \* H, halogen, alkyl, alkoxy, aryloxy, acylamino; X5 \* a bond or CMe2) can be used in rapid processable color photog, materials to give yellow images with outstanding grain and no color fog resulting from the presence of a fixing agent in the color developers. Developers for use with these materials contain 20.029 mol/L of Br. Thus, a gelatin Ag(Rf, I) emulsion containing IV at 2 \* 10-1 mol/mol Ag halided on a support at 1.2 g Ag/m2, dried, step wedge exposed, and color developed in a developer containing NaBr 7.0 g/L to give a RMS granularity of 41 vs. 76 for a control developed in a developer containing NaBr 1.2 g/L.

RL: USES (Uses)

(photog. yellow coupler for images with improved grain and decreased fog)

53812-46-7 CAPLUS

2-Benzofurancarboxamide, 5-(1,1-dimethylethyl)-N-[2-(hexadecyloxy)-5-[(methylamino)sulfonyl]phenyl]-2,3-dihydro-3-oxo- (9CI) (CA INDEX NAME)

Robert Haylin 107:113476 2,3-Dihydrobenzo[b]furan derivatives Boeckelmann, Juergen; Fanghaenel, Egon; Grossmann, INVENTOR (S): Norbert VEB Filmfabrik Wolfen, Fotochemisches Kombinat, Ger. PATENT ASSIGNEE(S): Dem. Rep. Ger. (East), 4 pp. CODEN: GEXXAS SOURCE: DOCUMENT TYPE: Patent FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO DATE DD 237164
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
GI DD 1985-276162 DD 1985-276162 A1 19860702 19850510 CASREACT 107:115478

Benzofurans I [R1, R2 = H, halo, NO2, carbalkoxy, acylamido, R4, R5 = (un)substituted alkyl, cycloalkyl, or aryl, alkoxy, aryloxy, (un)substituted NH2], useful as intermediates for plant protective agents, pharmaceuticals, aromas, and in the photog. industry, were prepared by, cyclization of phenols II [X = halo, 0502R3, R3 = (un)substituted aryl] with R4COCHYCOSS (Y = halo) via an intermediate betaine. 2,4-clcH2(02N)C6H3OH in Me2CO was treated with NEt3 in Me2CO to give 85V intermediate betaine which cyclized with MeCOCHCCO2Et and NEt3 in refluxing MeCN to give intermediate betaine which cyclized with MeCOCHCCO2Et and NEt3 in refluxing MeCN to give .apprx.90% I [R1 = NO2, R2 = H, R4 = OET, R5 = Me].

110110-75-3F

RL: SPN (Synthetic preparation), PREP (Preparation)
(preparation of, as intermediate for photog. substances, pharmaceuticals, plant protectants, and aromas)
110110-75-3 CAPLUS
2-Benzofurancarboxamide, 2-(2,2-dimethyl-1-oxopropyl)-2,3-dihydro-5-nitro-N-phenyl- (9CI) (CA INDEX NAME)

Robert Havlin

ANSWER 17 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN SSION NUMBER: 1979:481551 CAPLUS <u>Full-text</u>

10/553,108

53812-50-3 CAPLUS

Benzofurancarboxamide, 5-(1,1-dimethylethyl)-2,3-dihydro-N-[5-methyl-2-pctadecyloxy)phenyl]-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 18 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STW
ACCESSION NUMBER: 1975:9972 CAPLUS Full-text
BOCUMENT NUMBER: 2:9972
FITTLE: Photographic two-equivalent yellow couplers
Kunitz, Priedrich W., Kirchhoff, Gertrud
Agfa-Gevaert A.-G.
Ger Offen., 17 pp.
CODEN: GMXXEX

DOCUMENT TYPE:

DOCUMENT TYPE: LANGUAGE: Patent German FAMILY ACC. NUM. CO PATENT INFORMATION:

> PATENT NO. KIND DATE APPLICATION NO. DATE DE 1973-2313989 BE 1974-1005793 CA 1974-195423 GB 1974-12564 19740926 19740916 19730321 DE 2313989 BE 812283 19740314 CA 1016385 19740319 19760505

RL: USES (Uses)

(photog. 2-equivalent yellow coupler) 53812-46-7 CAPLUS

23-Benzofurancarboxamide, 5-(1,1-dimethylethyl)-N-[2-(hexadecyloxy)-5-(methylamino)sulfonyl]phenyl]-2,3-dihydro-3-oxo- (9CI) (CA INDEX NA

53812-47-8 CAPLUS 2-Benzofurancarboxanide, N-[5-[(dodecylamino)carbony1]-2-methoxypheny1]-2,3-dihydro-5-methyl-3-oxo- (SCI) (CA INDEX NAME)

53812-48-9 CAPLUS

2-Benzofurancarboxamide, N-[2-(diethylamino)-5-((octadecylamino)sulfonyl)phenyl]-2,3-dihydro-5-methyl-3-oxo- (9CI) (CA INDEX NAME)

osel2-49-0 CAPUOS 2-Benzofurancarboxamide, N-{2-{hexadecyloxy}phenyl}-2,3-dihydro-5-methyl-3-oxo- (9CI) (CA INDEX NAME)

10/553,108

39/40

Robert Havlin

L4 ANSWER 20 OP 20 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 1963:403485 CAPLUS Pull-text
DOCUMENT NUMBER: 59:3485

59:606b-h.607a-b ORIGINAL REFERENCE NO. :

TITLE: 2-Formyl-1,4-benzodioxane and 2-formyl-2,3-dihydrobenzofuran

AUTHOR (S)

Waisti, Domenico, De Marchi, Franco, Rosnati, Vittorio Ist. Super. Santta, Rome Gazzetta Chimica Italiana (1963), 93, 52-64 CODEN SOCITAS, ISSN. 0016-5603

CORPORATE SOURCE: DOCUMENT TYPE:

Journal Unavailable

CODEN. GCITAP, ISSN: 0016-5603

UNENT TYPE:

CODEN. GCITAP, ISSN: 0016-5603

UNENT TYPE:

JOAVAILABILE

For diagram(s), see printed CA Issue.

cf. CA 57, 8561h. PNNHME (0.17 mole) in 100 ml. dry CHCl3 slowly treated with 0.075 mole

1.4-benzodioxane-2-carboxylic acid chloride in 50 ml. CHCl3 and the mixture refluxed 4

hrs., washed with dilute HCl, aqueous Na2col, and H20, and the dried (Cacl2) solution

evaporated, the residue distilled at 160-59/0.08 mm., and the oily product (20 g.)

recrystd. from dilute MCDH gave the anilide (I, R = COMMeth) (II), m. 91-2°. II (47.5

g.) in 250 ml. dry tetrahydrofuran stirred at -5° with addition of 220 ml. 0.45M LiAlH4

in tetrahydrofuran below 0° and the mixture stirred 2 hrs. at 7°, treated with 360 ml. 9N

H2504, the acid aqueous layer diluted with 300 ml. H2O, saturated with NaCl, and

extracted repeatedly with Et2O, the dried (Cacl2) extract evaporated, and the residue

distilled yielded 60° I (R = CHO) (III), b0.08 69°-72°, transformed on long standing into

a glassy substance, (C9H803)1.8120, m. 64-7°, and 7.5 g, partially crystalline fraction,

b0.05, 85-95° recrystd. from 1:1 C6H6-C6H14 to give 1.5 g. I (R = CHOM) (IV), m. 84-6°.

The mother liquors concentrated and the residue distilled gave 6 g, hemiacetal 0f III

with IV), b0.04 90-5° v 3448, 3125, 2899, 1595, 1497, 1464, 1991, 1302, 1269, 1198, 1147,

110-1040, 927, 905, 840,750 cm.-1 (neat), also prepared by refluxing equimolar ants. of

III and IV in CHCl3 saturated with dry HCl. The distillation gave a 3rd viscous liquid

fraction (5 g.), b0.05 140-200°, infrared spectrum in ChCl3 lacking absorption bands in

the Co region and showing a smooth badly resolved band in the 1200-800 cm.-1 region.

PNNHMe (10, 110-mole) in 30 m. dry C6H6 added with stirring and cooling (ice bands and the product crystallized from Et2O yielded 96% I (R =

CH2CONNEPH) (V), m. 85-7°, recrystd. to give a sample, m. 87-8° (C6H14). V (0.025 mole)

in 250 ml. Et2O at -5° stirred with gradual addition of 25 ml. 0.55 m LiAlH4 in Et2O

(

10/553,108

2 Benzofurancarboxamide, 5-(1,1-dimethylethyl)-2,3-dihydro-N-[5-methyl-2-(octadecyloxy)phenyl]-3-oxo-(9CI) (CA INDEX NAME)

53812-51-4 CAPLUS

2-Benzofurancarboxamide, N-{2-chloro-4-(hexadecyloxy)pheny1}-2,3-dihydro-5-methyl-3-oxo-(9CI) (CA INDEX NAME)

ANSWER 19 OF 20 CAPLUS COPYRIGHT 2007 ACS ON STN SSION NUMBER: 1974:108307 CAPLUS Full-text MENT NUMBER: 80:108307

ACCESSION NUMBER: DOCUMENT NUMBER:

ACCESSION NUMBER: 80:108307

TITLE: Synthesis of abutic acid (5,6-dimethoxycoumarons-2,3-dicarboxylic acid)

AUTHOR(S): Jha, O. P.

CORPORATE SOURCE: Dep. Chem., Bhagalpur Univ., Bhagalpur, India

SOURCE: Indian Journal of Chemistry (1973), 11(10), 989-90

CODEN: IJOCAP, ISSN: 0019-5103

DOCUMENT TYPE: Journal

LANGUAGE: Bnglish

CI For diagram(s), see printed CA Issue.

AB 2,4,5-(MeO)3C6H2CO2H was treated with SOC12 and CH2N2 and the resulting diazoacetophenone cyclized with HoAc to give 5,6-dimethoxy-2-hydroxyphenyl)glyoxylic acid. The glyoxylic acid was cyclized with Aco2 and the resulting 5,6-dimethoxy-2-hydroxyphenyl)glyoxylic acid. The glyoxylic acid was cyclized with Aco2 and the resulting 5,6-dimethoxy-2-dione cleaved with MeONa to give Me [2-(methoxycarbonyl)methoxy-4,5-dimethoxyphenyl]glyoxylate which was cyclized with MeONa to give abutic acid 1.

52196-52-8P EATHOR-SALEP
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
52196-52-8 CAPLUS

2.3-Benzofurandicarboxamide, 2,3-dihydro-5,6-dimethoxy-N,N'-diphenyl-(9CI) (CA INDEX NAME)

A0/40

Added at 0\* with stirring to 0.113 mole 2,3-dihydrobenzofuran-2-carboxylic acid chloride in 100 ml. Et20, the mixture kept 15 hrs., the Et20 evaporated, and the diazo ketone taken up in 250 ml. absolute alc., stirred at 50\* with 2.4 g. Ag20 in 10 ml. absolute alc., and the mixture refluxed 8 hrs., the filtered molution evaporated, and the oily residue distilled gave 9 g. VII (R = .CH2CO2Et) (XII), bb. 05 100-20\*, and 8 g. yellow oil, bb. 03 125-8\* discarded). XII (S. S. g.) in 30 ml. alc. refluxed 1 hr. with 15 ml. 2N NaON and the cooled hydrolyzate extracted repeatedly with Et20, the aqueous alkaline solution acidified and exhaustively extracted with Et20, the dried extract evaporated, and the residue distilled yielded 4 g. XI, bb. 06 140-50\*, m. 76-9\* (CSH14), probably a dimorphic crystalline variant of the above XI. XI (3 g.) in 20 ml. C6He refluxed 45 min. with 3 ml. SOC12 and the residue on evaporation distilled yielded 3 g. VII (R = CH2COC1), bb.1 90-2\*, converted by treatment with PhNMePh in C6He to yield 92\* product, recrystd. from C6He-C6H4 to give VII (R = CH2COMPh) (XIII), m. 95-7\*. XIII (0.011 mole) in 150 ml. dry Bt20 at -6\* stirred with gradual addition of 13 ml. 0.53 ml.LAIH4 (0.007 mole) and the mixture kept 7 hrs. at -5\*, treated with 25 ml. SN H2SO4, and the aqueous phase extracted twice with Et20, the dried (Na2SO4) extract evaporated, and the oily residue distilled yielded 85\* VII (R = CH2CHO), bb.01 84-6\*, v 3030, 2899, 2817, 2703, 1727, 1597, 1481, 1462, 1377, 1325, 1225, 1227, 117, 1036, 1016, 983, 922, 870, 794, 750, 710 cm.-1 (neat)) p-nitrophenylhydrasone m. 146-50\* (decomposition) (alc.). Comparison of the properties of the 2 aldehydes III and X with compds. of closely related structure polymeric aldehydes. 92362-25-10, 2-8 enzofurancarboxanilide, 2,3-dihydro-N-methyl- (preparation of) 92962-85-1 (CAPUS

(preparation of 92962-85-1 CAPLUS

2-Benzofurancarboxanilide, 2,3-dihydro-N-methyl-.(7CI) (CA INDEX NAME)

c*>		
-> log hold		
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